# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

		<u></u> *
(51) International Patent Classification 7:		(11) International Publication Number: WO 00/54823
A61M 1/36, 1/02, B04B 13/00	A1	(43) International Publication Date: 21 September 2000 (21.09.00)
(21) International Application Number: PCT/USC		61 (81) Designated States: AU, CA, JP, European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC,
(22) International Filing Date: 14 March 2000 (	14.03.0	

US

(71) Applicant: TRANSFUSION TECHNOLOGIES CORP.
[US/US]; 9 Erie Drive, Natick, MA 01760 (US).

17 March 1999 (17.03.99)

(72) Inventor: HEADLEY, Thomas, D.; 83 Westgate Road, Wellesley, MA 02181 (US).

(74) Agents: SUNSTEIN, Bruce, D. et al.; Bromberg & Sunstein LLP, 125 Summer Street, Boston, MA 02110-1618 (US).

Published

With international search report.

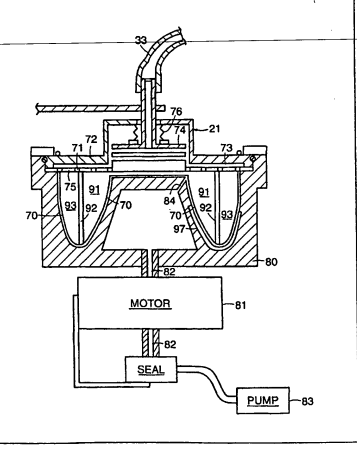
(54) Title: SYSTEM AND METHOD FOR COLLECTING PLATELETS AND OTHER BLOOD COMPONENTS

#### (57) Abstract

(30) Priority Data:

09/271,601

A system for collecting, from whole blood, platelets suspended in plasma. By centrifuging the blood at a high enough rotational speed, the platelets (92) are separated from the plasma (91) and the red blood cells (93). In a preferred embodiment, some of the plasma (91) is removed while the centrifuge is being spun to keep the platelets (92) separated from the plasma (91). Then, the speed of rotation is altered so as to cause the platelets (92) to mix with the remaining plasma (91). The platelets (92) can then be collected with the remaining plasma (91).



#### FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

İ	AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
i	AM	Аплеліа	Fi	Finland	LT	Lithuania	SK	Slovakia
I	AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
١	ΑU	Australia	GA	Gabon ´	LV	Latvia	SZ	Swaziland
۱	ΑZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
ı	BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
ł	ВВ	Barbados	GH	Ghana	MG-	Madagascar	TJ	Tajikistan
l	BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
1	BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
l	BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
١	BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
1	BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
	BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
ı	CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
	CF	Central African Republic	JР	Japan	NE	Niger	VN	Viet Nam
	CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
ĺ	CH	Switzerland	KG	Kyrgyzstan	NO	Norway	2W	Zimbabwe
	CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
	СМ	Cameroon		Republic of Korea	PL.	Poland		
ļ	CN	China	KR	Republic of Korea	PT.	Portugal		
	cυ	Cuba	KZ	Kazakstan	RO	Romania		
	cz	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
	DE	Germany	LI	Liechtenstein	SD	Sudan		
	DK	Denmark	LK	Sri Lanka	SE	Sweden		
	EE	Estonia	LR	Liberia	SG	Singapore		

# SYSTEM AND METHOD FOR COLLECTING PLATELETS AND OTHER BLOOD COMPONENTS

5

#### Technical Field

This invention generally relates to systems and methods for processing blood and other biological fluids.

10

#### Background Art

Fig. 1 shows a typical disposable bag set used in the prior art to collect platelets from whole blood. The set includes a needle 10 or cannula, which is inserted into a vein of a donor. The needle 10 is connected to the tube 11, which in turn is connected to collection bag 12, so as to allow whole blood to flow from the donor through the needle 10 and the tube 11 into collection bag 12. The collection bag 12 contains anticoagulant. After the desired amount of blood has been collected into collection bag 12, the needle 10 is removed from the donor, and tube 11 is cut and heat sealed. The remainder of the bag set is then brought to a centrifuge, which spins the bag set so that the blood in collection bag 12 separates into platelet-rich plasma and red blood cells. Typically, the centrifuge is not located at the point where the blood donation takes place.

After the blood has separated into platelet-rich plasma and red blood cells (RBCs), the bag set is removed from the centrifuge. The platelet-rich plasma is urged from collection bag 12 through tube 13 into platelet-storage bag 14. The tube 13 leading to the platelet- and plasma-storage bags 14, 15 is then cut and heat sealed. Storage-solution bag 16 holds RBC-storage solution. After the platelet-rich plasma has been urged-into the platelet-storage bag 14, the RBC-storage solution is urged from the storage-solution bag 16 into the collection bag 12. The tube 41 connecting the collection and storage-solution bags 12, 16 is then cut and heat sealed.

At this stage, the bag set has been divided into two portions: (i) the first portion consists of the collection bag 12, which now holds primarily red blood cells (along with storage solution), filter 17, RBC-storage bag 18, and the tubing 19 that connects these components, and (ii) the second portion consists of the platelet-storage bag 14, which now holds platelet-rich plasma, and the plasmastorage bag 15 and the tubing that connects these two components.

The first portion may be hung, so that gravity causes the RBC component to pass from the collection bag 12 through the filter 17 to RBC-storage bag 18.

The filter 17 removes white blood cells (WBCs) from the red blood cells. After the red blood cells (and storage solution) pass into the RBC-storage bag 18, tube 19 is cut and heat sealed.

To collect platelets, the second portion is centrifuged at a high rotational speed in order to separate the platelets from the plasma. After the platelets have been separated from the plasma, some of the plasma is urged from the platelet-storage bag 14 into the plasma-storage bag 15. Typically, 50 mls of plasma are left with the platelets in the platelet-storage bag 14. After the desired amount of plasma has been removed from the platelet-storage bag 14 to the plasma-storage bag 15, the tube connecting these two bags is cut and heat sealed. Thus, at the end of the procedure, the platelet-storage bag 14 holds platelets in about 50ml of plasma, the plasma-storage bag 15 holds platelet-poor plasma, and the RBC-storage bag 18, of course, holds red blood cells.

This prior-art process of collecting and separating blood components involves many steps and frequent human intervention. The arrangement of the prior-art bag set does not permit the process to be easily automated.

Summary of the Invention

25

The present invention is directed to systems and methods for collecting, from whole blood, platelets suspended in plasma. By centrifuging the blood at a high enough rotational speed, the platelets are separated from the plasma and the red blood cells. In a preferred embodiment, some of the plasma is removed

while the centrifuge is being spun to keep the platelets separated from the plasma. Then, the speed of rotation is altered so as to cause the platelets to mix with the remaining plasma. The platelets can then be collected with the remaining plasma.

5

A system that may be used for carrying out the invention includes a centrifuge rotor, a flow-control arrangement and a spinner. The flow-control arrangement introduces whole blood into the centrifuge rotor and removes blood components from the centrifuge rotor. A controller causes the spinner to rotate at two different speeds: The rotor is spun at a first speed so as to separate 10 the blood into a first component, a second component and a third component. The first component is primarily plasma. The second component is located, while the rotor is being spun, outside of the first component and is primarily red blood cells. The third component is located, while the rotor is being spun, between the first and second components and includes platelets. The controller 15 causes the rotor's speed of rotation to be altered so as to cause the third component to mix with the first component. The controller also causes the flowcontrol arrangement to remove from the rotor a portion of the plasma containing platelets.

As noted above, in a preferred embodiment, the controller causes the 20 flow-control arrangement to remove some of the first component (the plasma) before the third component (comprising the platelets) is mixed with the first component. The system also preferably includes a plasma-volume determination sensor in communication with the controller; the plasma-volume determination sensor determines the volume of the first component in the rotor.

25 The controller may thus remove a portion of the first component based on the determined volume of the first component.

### Brief Description of the Drawings

10

FIG. 1 shows a disposable set that may be used in a prior-art system for collecting platelets from whole blood.

FIG. 2 shows a cross-sectional view of a variable-volume rotor mounted in a chuck that spins the rotor and causes the rotor's volume to change.

FIG. 3 shows a disposable set using a variable-volume rotor, such as the one in FIG. 2.

FIG. 4 shows a control unit holding the disposable of FIG. 3.

#### Description of Specific Embodiments

A method of collecting platelets is described in connection with FIG. 2. FIG. 2 shows a cross-section of a rotor 21 mounted in a chuck 80, which is located in the control unit and which holds the rotor 21. This rotor 21 may be any one of a variety of designs, but preferably the rotor has a variable total volume, such as the rotors shown and described in U.S. Patent No. 5,733,253 (which is incorporated herein by reference). (The rotor shown in FIG. 2 is similar to the rotor shown in Figs. 1-4 of U.S. Patent No. 5,733,253, but it will be appreciated that other designs, such as other designs shown in U.S. Patent No. 5,733,253, may be used instead.) A motor 81 causes the chuck 80 and the rotor

- 20 21 to spin. The control unit also includes a pump 83, which is connected through the cannulated axis 82 of the motor 81 to the interior of the chuck 80. The rotor 21 has an elastic diaphragm 70, which defines the interior volume of the rotor 21. Upper boundary wall 72 also defines the interior volume of the rotor 21. The position of the diaphragm 70 determines the volume of the rotor,
- and the position of the diaphragm 70 may be controlled by controlling, by means of the pump 83, the pressure of the gas in the interior of the chuck 80.

  The interior of the chuck 80 includes one or more apertures 84 to permit the gas to come into fluid communication with the diaphragm 70. The rotor 21 may also include a interior wall 75 with perforations 71. The boundary wall 72 and
- 30 the interior wall 75 form a passage 73, through which blood and blood

components may flow to and from the rotor's non-rotating portion 74 and the tubing 33 attached to the rest of the disposable set. A rotary seal 76 provides a seal between the rotating and non-rotating portions of the rotor 21. In lieu of the perforated interior wall 75, channels may be located on the interior surface of the boundary wall 72 to provide fluid communication between the rotor's non-rotating portion 74 and the outer radius of the rotor's interior (as shown in Figures 41 and 42 of above-referenced U.S. Patent No. 5,733,253).

FIG. 2 shows the rotor 21 at its maximum volume, with the diaphragm 70 stretched as far as the chuck 80 permits it to be stretched. The rotor 21 is spun sufficiently fast by the chuck 80 and the motor 81 to cause the blood to be separated into red blood cells 93, platelets 92 and plasma 91. Since, of these three blood components, the RBC component 93 of the blood has the greatest specific gravity, the RBC component is the furthest from the rotor's axis of rotation. The plasma component 91 has the lightest specific gravity, and therefore the plasma component is the closest to the axis of rotation. The platelet component 92, having an intermediate specific gravity, forms a thin layer between the plasma and RBC components.

In order to collect the platelets, it is preferable first to collect all but about 50 milliliters of the plasma. The remaining 50 mls of plasma will be used to store the platelets, as the standard practice in the industry is to store a unit of platelets in 50 mls of plasma. The plasma 91 is collected (i.e., urged through fixed portion 74 to tube 33) by continuing to spin the rotor 21 and using the pump 83 to increase the pressure against the diaphragm 70, and/or by slowing the revolutions of the rotor 21. The rotor 21 should preferably continue to be spun quickly and smoothly enough to keep the platelets 92 in a separate layer.

Once all but 50 mls of plasma 91 has been collected, the platelets 92 may be mixed with the remaining plasma by sharply changing the speed of rotation of the rotor 21. It has been found that, by sharply changing the rotor's speed of rotation, the platelets will mix with the neighboring plasma. Because the red blood cells have a much heavier specific gravity, the red blood cells tend to

remain in their separate layer. Of course, the rotor's speed must not be altered so radically and quickly as to cause the red blood cells as well to mix with the other components. Alternatively, the speed of rotation may be slowed sufficiently--although not necessarily sharply--so that the platelets mix with the plasma but the red blood cells remain separate. Once the platelets are mixed with the remaining plasma, additional pressure may be created by the pump 83 to push the diaphragm 70 further outward and force the platelets, now suspended in plasma, out of the rotor into tube 33. The red blood cells may then be collected. Each of the components, platelets suspended in plasma, platelet-poor plasma, and the red blood cells should be directed to a separate container. Alternatively, one or both of the platelet-poor plasma and the red blood cells may be returned to the donor.

FIG. 3 shows a disposable set that may be used in the platelet-collection process just described. The disposable set includes the rotor 21, a plasma-15 storage container 24, a platelet-storage container 99, a RBC-storage container 28, a filter 17 for removing white blood cells from the red blood cells, a cannula 10 (or other means for permitting whole blood to enter the disposable set), and tubing 33 connecting these components. The plasma-storage container 24 may contain anticoagulant, which may be introduced into the whole blood as it is 20 being drawn through the needle 10 to the rotor 21. The platelet-storage container 99 may contain platelet-storage solution, and the RBC-storage container 28 may contain RBC preservative. After the plasma and the platelets have been removed from the rotor 21, the RBC preservative may be urged from the RBC-storage container 28 into the rotor 21, where the RBC preservative is 25 mixed with the red blood cells remaining in the rotor 21. The red blood cells and the preservative may then be urged from the rotor 21 through the filter 17 into the RBC-storage container 28, in the manner described in concurrently filed application, serial no. \_\_/\_\_\_\_, for a "System and Method for Separating Blood Components," bearing attorney docket number 1611/112 and listing Headley and Powers as inventors. (This application is incorporated herein by

reference.) Alternatively, the present invention may be used with the system and process described in concurrently filed application, serial no. \_\_/\_\_\_\_, for a "System and Method for Red-Blood-Cell Apheresis," bearing attorney docket number 1611/119 and listing James Cianci as the inventor. (This application is also incorporated herein by reference.)

FIG. 4 shows the disposable set of FIG. 3 mounted in a control unit 20.

The control unit 20 includes a flow-control arrangement for controlling and/or causing flow between the needle 10, the rotor 21 and the storage containers 24, 99, 28. The flow-control arrangement may include valves 22, 23, 98, 27, which control the flow through the various branches of the tubing. Alternatively, a single valving cassette may be used to control the flow through the various branches of the tubing. By varying the pressure against the rotor's diaphragm (item 70 in FIG. 9), and by varying the speed that the rotor is spun 21, fluid may be urged into or out of the rotor from and to the needle 10 or the storage containers 24, 99, 28. For instance, applying a vacuum on the rotor's diaphragm while valve 22 is open helps draw blood from the donor into the rotor 21. In addition to or in lieu of changing the pressure against the rotor's diaphragm, the control unit may be provided with independent pumping mechanisms (such as a peristaltic pump) that act on the tubing (or on a valving cassette) to force fluid

a peristaltic pump) that act on the tubing (or on a valving cassette) to force fluid
through the tubing in the desired direction.

In order to determine how much plasma should be removed in order to

In order to determine how much plasma should be removed in order to leave only 50 mls of plasma, in which the platelets are to be suspended, the control unit may be provided with an arrangement for determining the volume of the red blood cells. One means of determining the volume of the red blood cells is to provide an array 97 of optical sensors (shown in FIG. 2) in the chuck 80 to determine the radius of the inner boundary of the red blood cells 93 when the blood has been centrifuged into different components. (If the boundary wall 72 is translucent, the array may be mounted above the rotor 21 instead of below it.) The control unit 20 may then calculate the volume of the red blood cells based on the location of this boundary when the rotor is filled with, say, one unit of

blood. Using this volume information, the control unit may determine approximately the weight of the red blood cells in the rotor, based on the specific gravity of red blood cells.

By weighing the chuck/rotor combination before and after the whole

blood was introduced into the rotor, the control unit may determine the weight of all the blood components in the rotor when the rotor is filled. By subtracting the weight of the red blood cells from the total weight of all the blood components in the rotor, the control unit may determine approximately the weight of the plasma in the rotor, and how much of it should be removed in

order to leave approximately 50 mls of plasma in the rotor. By weighing the chuck/rotor combination as platelet-poor plasma is being urged from the rotor, or alternatively by weighing the container 24 that holds the plasma as it leaves the rotor, the control unit can stop removing plasma when the correct amount of plasma has been removed. The platelet-poor plasma is preferably directed to

At that point, there should be approximately 50 mls of plasma left in the rotor, as well as all the platelets and all the red blood cells. The speed of the rotor may then be changed rapidly, in order to cause the platelets 92 to become mixed in the approximately 50 mls of the plasma remaining. The

20 platelet/plasma combination is then urged from the rotor and sent to the

- platelet/plasma combination is then urged from the rotor and sent to the platelet-storage container 99. Another optical sensor 96, mounted on the outlet tube 33 senses when the red blood cells start emerging from the rotor. (See FIG.
  - 4.) When the red blood cells are detected, flow to the platelet-collection container 99 is stopped, and the red blood cells may be directed through filter 17 to a RBC-collection container 28.

Although the invention has been described with reference to several preferred embodiments, it will be understood by one of ordinary skill in the art that various modifications can be made without departing from the spirit and the scope of the invention, as set forth in the claims hereinbelow.

What is claimed is:

1. A system of collecting a fluid component of intermediate specific gravity, mixed with a fluid component of lighter specific gravity, the system comprising: a centrifuge rotor;

a flow-control arrangement which introduces whole fluid into the centrifuge rotor and removes fluid components from the centrifuge rotor;

a spinner which rotates the rotor at more than one speed;

a controller for causing the spinner to rotate the rotor at a first speed so as to separate the fluid into a first component, a second component and a third component, wherein the first component has the lightest specific gravity, wherein the second component is located, while the rotor is being spun, outside of the first component and has the greatest specific gravity, and wherein the third component is located, while the rotor is being spun, between the first and second components and has an intermediate specific gravity, and for causing the rotor's speed of rotation to be altered so as to cause the third component to mix with the first component, and for causing the flow-control arrangement to remove from the rotor a portion of the first component containing the second component.

20 2. A system of collecting platelets suspended in plasma, the system comprising:

a centrifuge rotor;

25

a flow-control arrangement which introduces whole blood into the centrifuge rotor and removes blood components from the centrifuge rotor;

a spinner which rotates the rotor at more than one speed;

a controller for causing the spinner to rotate the rotor at a first speed so as to separate the blood into a first component, a second component and a third component, wherein the first component is primarily plasma, wherein the second component is located, while the rotor is being spun, outside of the first component and is primarily red blood cells, and wherein the third component is

located, while the rotor is being spun, between the first and second components and includes platelets, and for causing the rotor's speed of rotation to be altered so as to cause the third component to mix with the first component, and for causing the flow-control arrangement to remove from the rotor a portion of the plasma containing platelets.

3. The system according to claim 2, wherein the controller causes the flow-control arrangement to remove some of the first component before the third component is mixed with the first component.

10

- 4. The system according to claim 3, further including a plasma-volume determination sensor in communication with the controller, wherein the plasma-volume determination sensor determines the volume of the first component in the rotor, and wherein the controller removes a portion of the first component based on the determined volume of the first component.
  - 5. The system according to claim 2, wherein the centrifuge rotor has a variable total volume.
- 20 6. The system according to claim 5, wherein the centrifuge rotor is provided with a flexible diaphragm which defines the volume of the rotor.
- The system according to claim 6, further including a pump which varies air pressure adjacent the flexible diaphragm so as to vary the total volume of the
   centrifuge rotor.
  - 8. The system according to claim 7, wherein the centrifuge rotor is provided with a fixed portion, a rotatable portion and a rotary seal providing a seal between the fixed and rotatable portions.

9. A system of collecting platelets suspended in plasma, the system comprising:

a centrifuge rotor;

means for introducing whole blood into the centrifuge rotor and removing blood components from the centrifuge rotor;

means for (i) spinning the rotor at a first speed so as to separate the blood into a first component, a second component and a third component, wherein the first component is primarily plasma, wherein the second component is located, while the rotor is being spun, outside of the first component and is primarily red blood cells, and wherein the third component is located, while the rotor is being spun, between the first and second components and includes platelets, and (ii) for altering the rotor's speed of rotation so as to cause the third component to mix with the first component.

- 15 10. The system according to claim 9, further including a controller for causing the introducing means to remove from the rotor a portion of the first component containing platelets before the rotor's speed of rotation is altered.
  - 11. The system according to claim 10, further including means for
- 20 determining how much plasma should be removed from the rotor before the rotor's speed is altered.
  - 12. A system for collecting platelets from whole blood, the method comprising:
- 25 a disposable set having

an inlet,

a platelet container,

a centrifuge rotor, and

tubing connecting the inlet, the platelet container and the rotor;

30 a control unit having

a spinner in which the rotor may be held and spun, and a controller for causing the spinner to rotate the rotor so as to separate the whole blood into a first component, a second component and a third component, wherein the first component is primarily plasma, wherein the second component is located, while the rotor is being spun, outside of the first component and is primarily red blood cells, and wherein the third component is located, while the rotor is being spun, between the first and second components and includes platelets, and then altering the rotor's speed of rotation so as to cause the third component to mix with the first component.

13. The system according to claim 12, wherein the control unit further includes a flow-control arrangement, which urges a mixture of the first and third components out of the rotor, while the rotor is still spinning.

15

10

- 14. The system according to claim 12, wherein the flow-control arrangement causes a portion of the first component to be urged from the rotor before the third component is mixed with the first component.
- 20 15. The system according to claim 12, wherein the centrifuge rotor has a variable total volume.
  - 16. A method of collecting platelets, the method comprising the steps of: introducing whole blood into a centrifuge rotor;
- spinning the rotor at a first speed so as to separate the blood into a first component, a second component and a third component, wherein the first component is primarily plasma, wherein the second component is located, while the rotor is being spun, outside of the first component and is primarily red blood cells, and wherein the third component is located, while the rotor is being spun, between the first and second components and includes platelets;

changing the rotor's speed of rotation so as to cause the third component to mix with the first component; and

removing from the rotor a portion of the plasma containing platelets.

- 5 17. The method according to claim 16, further including the step of removing a portion of the plasma from the rotor before the rotor's speed is changed.
  - 18. A method of collecting platelets from whole blood, the method comprising:
- providing a disposable set having an inlet, a platelet container, a centrifuge rotor, and tubing connecting the inlet, the platelet container and the rotor;

providing a control unit having a spinner in which the rotor may be held; placing the rotor in the spinner;

- drawing whole blood through the inlet;

  directing the whole blood from the inlet through the tubing to the rotor;

  causing the spinner to rotate the rotor so as to separate the whole blood into a first component, a second component and a third component, wherein the first component is primarily plasma, wherein the second component is
- 20 located, while the rotor is being spun, outside of the first component and is primarily red blood cells, and wherein the third component is located, while the rotor is being spun, between the first and second components and includes platelets;

altering the rotor's speed of rotation so as to cause the third component to
25 mix with the first component; and

urging a mixture of the first and second components out of the rotor, while the rotor is still spinning.

19. The method according to claim 18, wherein a portion of the first component is urged from the rotor before the third component is mixed with the first component.

- 5 20. The method according to claim 19, wherein the centrifuge rotor has a variable total volume.
- 21. The method according to claim 20, wherein whole blood is drawn through the inlet and directed to the rotor while the rotor is mounted in thespinner.
- 22. The method according to claim 21, wherein the centrifuge rotor is provided with a fixed portion, a rotatable portion and a rotary seal providing a seal between the fixed and rotatable portions, and the tubing is connected to the rotor's fixed portion.
  - 23. The method according to claim 22, wherein the control unit varies the volume of the centrifuge rotor.
- 20 24. The method according to claim 23, wherein the centrifuge rotor is provided with a flexible diaphragm which defines the volume of the rotor, and the control unit is provided with means for varying air pressure adjacent the flexible diaphragm so as to vary the total volume of the centrifuge rotor.
- 25. The method according to claim 20, wherein the centrifuge rotor is provided with a fixed portion, a rotatable portion and a rotary seal providing a seal between the fixed and rotatable portions, and the tubing is connected to the rotor's fixed portion.

26. The method according to claim 19, wherein the disposable set includes a plasma-storage container and an RBC container, and wherein the method further includes the step of directing a portion of the first component to the plasma-storage container and the third component to the RBC container.

5

27. The method according to claim 19, wherein the disposable set includes return means, and wherein the method further includes the step of directing at least one of the first and second components back to the donor through the return means.

10

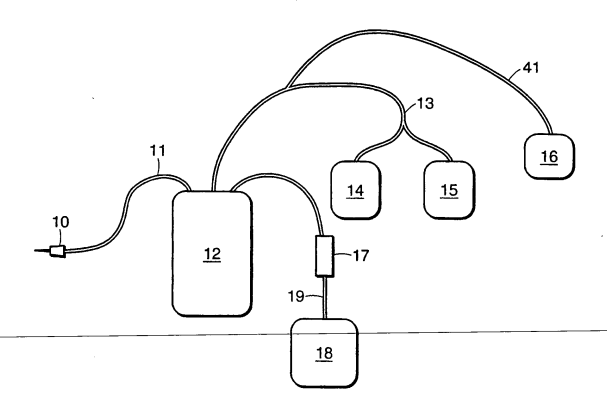


FIG. 1

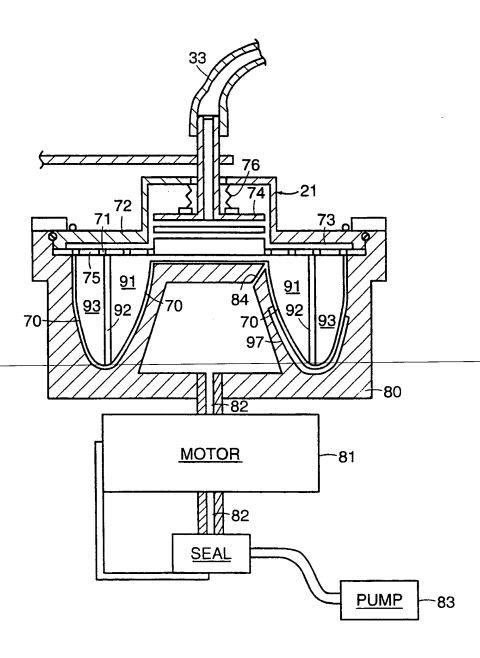


FIG. 2

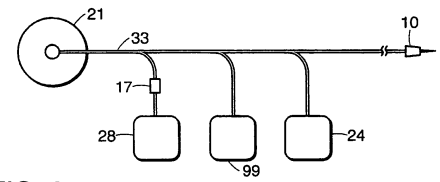
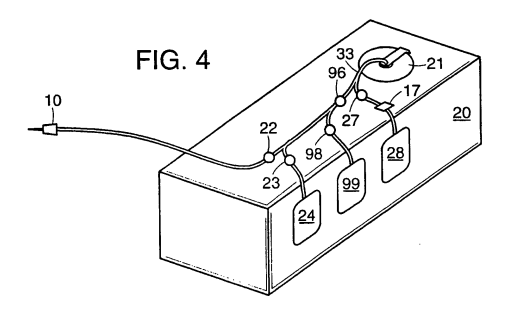


FIG. 3



# INTERNATIONAL SERCH REPORT

Inter onal Application No PCT/US 00/06561

			, 55 55, 60001
A CLASSI IPC 7	FICATION OF SUBJECT MATTER A61M1/36 A61M1/02 B04B13	3/00	
According to	o International Patent Classification (IPC) or to both national class	sification and IPC	
	SEARCHED		
Minimum do IPC 7	ocumentation searched (classification system followed by classifi A61M B04B	cation symbols)	
Documenta	tion searched other than minimum documentation to the extent the	at such documents are included in	the fields searched
Electronic d	ata base consulted during the international search (name of data	base and, where practical, search	i terms used)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
X	US 5 733 253 A (POWERS EDWARD 1 31 March 1998 (1998-03-31) cited in the application the whole document	ET AL)	1,2, 5-10,12, 15
A	in particular see: column 7, line 23 - line 45 column 11, line 56 -column 12, column 12, line 39 - line 43		11
A	column 21, line 9 -column 22, l figure 3	ine 14	16,18
A .	W0 96 33023 A (COBE LAB) 24 October 1996 (1996-10-24)		1,2,9, 12,16,18
	page 27, paragraphs 2,4; figure	9 5	
		-/	
X Furt	her documents are listed in the continuation of box C.	X Patent family membe	rs are listed in annex.
"A" docume consic "E" earlier e filling c "L" docume which citatio "O" docume other e "P" docume later ti	ategories of cited documents:  ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another or other special reason (as specified)  ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but han the priority date claimed	or priority date and not in cited to understand the privention  "X" document of particular relecannot be considered not involve an inventive step  "Y" document of particular relecannot be considered to it document is combined with	vel or cannot be considered to when the document is taken alone invance; the claimed invention involve an inventive step when the th one or more other such docubeing obvious to a person skilled same patent family
	June 2000	09/06/2000	massiai orangi lapurt
Name and I	mailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2  NL - 2280 HV Rijswijk  Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer  Lakkis, A	

# INTERNATIONAL SERCH REPORT

Inte. cnal Application No PCT/US 00/06561

	ation) DOCUMENTS CONSIDERED TO BE RELEVANT		Delevent to state to
ategory °	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
A	EP 0 885 619 A (TERUMO CORP) 23 December 1998 (1998-12-23) page 5, line 55 -page 6, line 1 page 8, line 56 - line 58		1
		•	

# INTERNATIONAL SERCH REPORT

information on patent family members

Inte conal Application No PCT/US 00/06561

Patent document cited in search report	ı	Publication date	1	Patent family member(s)		Publication date
US 5733253	Α	31-03-1998	AU	699141	В	26-11-1998
			AU	4132296	Α	06-05-1996
			CA	2202284	Α	25-04-1996
			EP	0839072	Α	06-05-1998
			JP	11511686	T	12-10-1999
			MO	9611747	Α	25-04-1996
			US	6039711	Α	21-03-2000
			US		Α	01-02-2000
			US	5885239		23-03-1999
			US	5904355	Α	18-05-1999
WO 9633023	Α	24-10-1996	US	5674173	Α	07-10-1997
			AU	702151	В	18-02-1999
			AU	5560896	Α	07-11-1996
			BR	9608031	Α	30-11-1999
			CA		Α	24-10-1996
			EP	1000664	Α	17-05-2000
			EP		Α	25-02-1998
			JP	11503966	T	06-04-1999
			US	5939319	Α	17-08-1999
			US		Α	03-03-1998
			US		Α	25-04-2000
			US		Α	08-02-2000
			US		A	22-06-1999
			US		Α	14-09-1999
~~~~~~~~			US	5906570	A	25-05-1999
EP 0885619	Α	23-12-1998	JP	11004886	A	12-01-1999